

Neurodegenerative disorders-1

Manar Hajeer, MD, FRCPath

University of Jordan, School of medicine

Classic features:

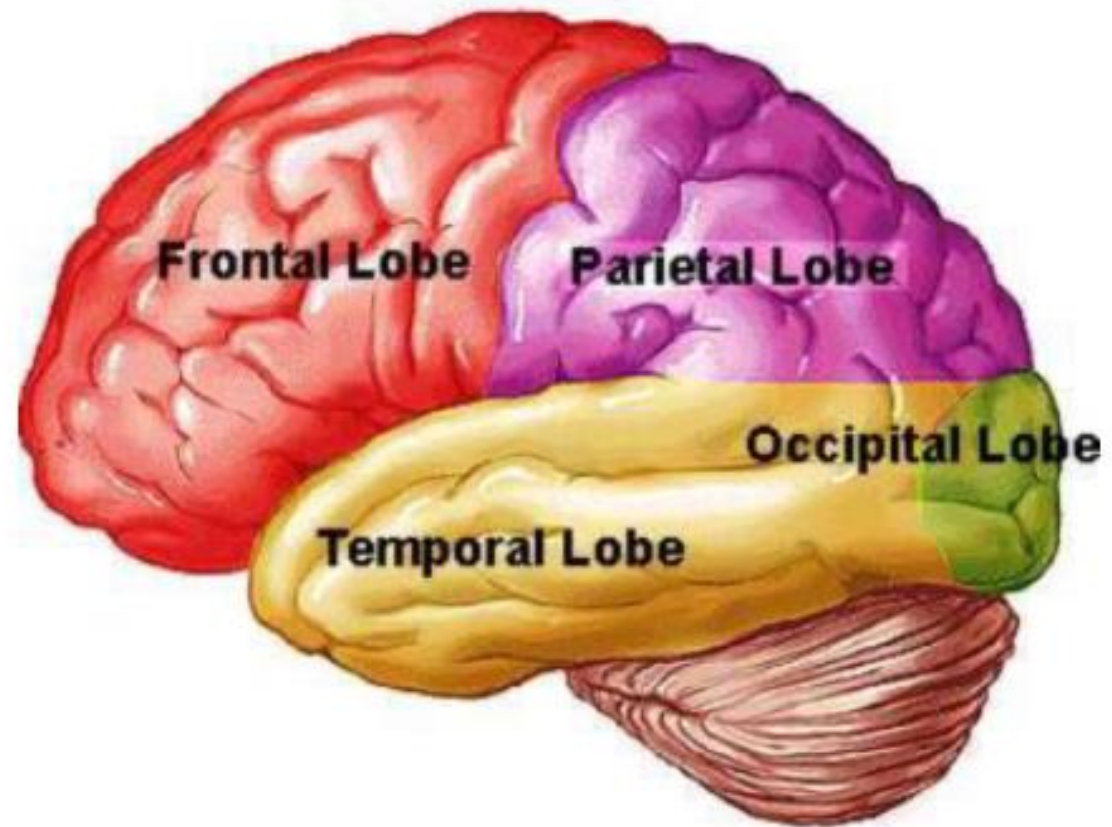
- ▶ Progressive loss of neurons.
- ▶ Typically affects groups of neurons with functional interconnections.
- ▶ Different diseases involve different neural systems, so different symptoms.
- ▶ The histologic hallmark for ALL diseases is the ACCUMULATION OF PROTEIN AGGREGATES.
- ▶ Same protein may aggregate in different diseases, BUT AT DIFFERENT DISTRIBUTION..
- ▶ Proteins resist degradation, accumulate within the cells, elicit inflammatory response, and is toxic to neurons.

Causes of protein accumulation

- ▶ Mutations that alter protein conformation.
- ▶ Mutations disrupting the processing and clearance of proteins.
- ▶ Subtle imbalance between protein synthesis and clearance (genetic or environmental factors)

Different diseases

- ▶ **Involving the hippocampus and cortex** >>>> cognitive changes (memory disturbances, behavior and language) >>>> dementia >>>> ALZHEIMER DISEASE (AD) , FRONTOTEMPORAL DEMENTIA (FTD), PICK DISEASE (SUBTYPE OF FTD)
- ▶ **Involving the basal ganglia** >>>> movement disorders >>>> hypokinesia (PARKINSON DISEASE) or hyperkinesia (HUNTINGTON DISEASE)
- ▶ **Involving the cerebellum** >>>> ataxia >>> (SPINOCEREBELLAR ATAXIA, FRIEDRICH ATAXIA, ATAXIA TELANGECTASIA)
- ▶ **Involving the motor system** >>> difficulty swallowing and respiration with muscle weakness >> (AMYOTROPHIC LATERAL SCLEROSIS)



Common features to many neurodegenerative diseases:

- ▶ Protein aggregates can seed the development of more aggregates.
- ▶ Protein aggregates can spread from one neuron to another in **Prion-like pattern**.
- ▶ No evidence of person-to-person transmission.
- ▶ Activation of the innate immune system is a common feature of neurodegenerative diseases.

DEMENTIA

- ▶ Development of *memory impairment* and other *cognitive deficits* severe enough to decrease the person's capacity to function at **his previous level** **despite** normal level of consciousness.
- ▶ Cognitive deficit *must affect the person's performance in his daily life activities.*
- ▶ There is no standard NORMAL COGNITION, always compared to previous level.

SYMPTOMS OF DEMENTIA

Cognitive changes

- Memory loss, which is usually noticed by a spouse or someone else

- Difficulty communicating or finding words

- Difficulty reasoning or problem-solving

- Difficulty handling complex tasks

- Difficulty with planning and organizing

- Difficulty with coordination and motor functions

- Confusion and disorientation

Psychological changes

- Personality changes

- Depression

- Anxiety

- Inappropriate behavior

- Paranoia

- Agitation

- Hallucinations

Causes of dementia

- ▶ Neurodegenerative diseases.
- ▶ Infections.
- ▶ Nutritional deficiencies.
- ▶ Metabolic and endocrine abnormalities
- ▶ Drugs.
- ▶ Subdural hematoma.
- ▶ Poisons.
- ▶ Tumours.
- ▶ Anoxia and ischemia.

COMPLICATIONS OF DEMENTIA

- ▶ **Inadequate nutrition.** Many people with dementia eventually reduce or stop their intake of nutrients.
- ▶ **Inability to perform self-care tasks.** As dementia progresses, it can interfere with bathing, dressing, brushing hair or teeth, using the toilet independently and taking medications accurately.
- ▶ **Personal safety challenges.** Some day-to-day situations can present safety issues for people with dementia, including driving, cooking and walking alone.
- ▶ **Death.** Late-stage dementia results in coma and death, often from infection

Alzheimer disease:

- ▶ Most common cause of dementia in older adults.
- ▶ Increase incidence with age (47% in those over 84 years).
- ▶ Most cases are sporadic.
- ▶ 5-10% are familial (onset before 50)
- ▶ Gradual onset.
- ▶ Impaired higher intellectual functions, memory impairment and altered mood and behavior.
- ▶ Severe cortical dysfunction with time (disorientation and aphasia, profound disability, mute and immobile)
- ▶ Death usually due to infections (pneumonia)

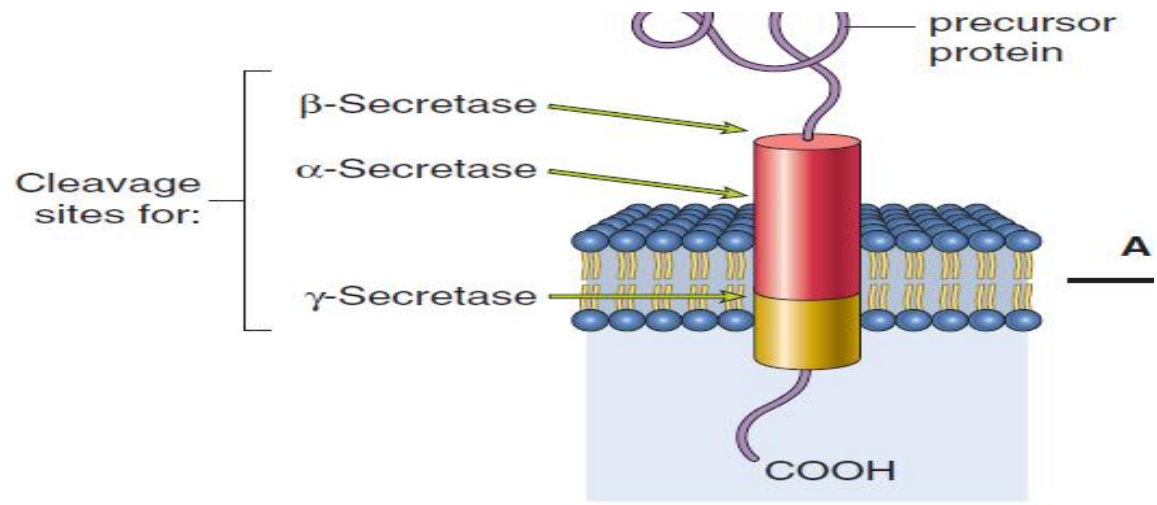
- ▶ **The most commonly recognized symptom of Alzheimer is an inability to acquire new memories and difficulty in recalling recently observed facts.**
- ▶ As the disease advances, symptoms include confusion, irritability and aggression, mood swings, language breakdown, long term memory loss, and ultimately a gradual loss of bodily functions and death.

Pathogenesis:

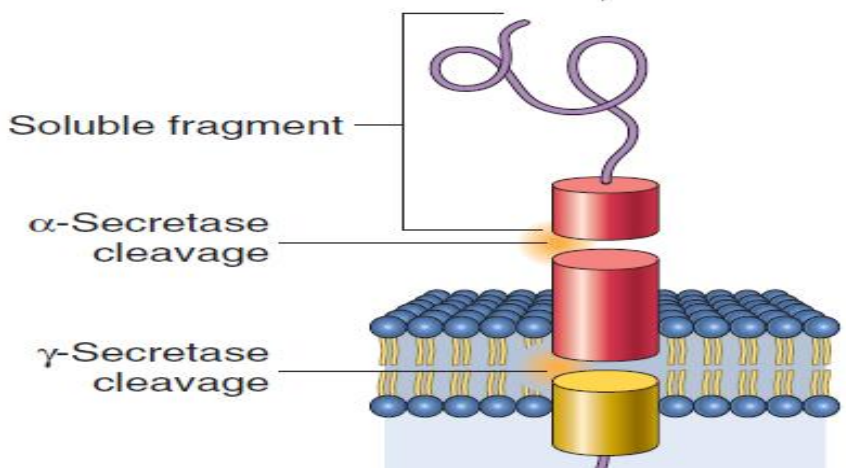
- ▶ Accumulation of two proteins (A β amyloid and Tau)
- ▶ In the form of plaques and neurofibrillary tangles, respectively.
- ▶ This leads to neuronal dysfunction, death and inflammation.
- ▶ Plaques deposit in the neuropil.
- ▶ Tangles develops intracellularly.
- ▶ A β generation is the critical initiating event for the development of AD.
- ▶ Mutations of the gene encoding the precursor protein for A β >>> elevated risk of AD.
- ▶ Mutations of Tau gene do NOT increase risk of AD.

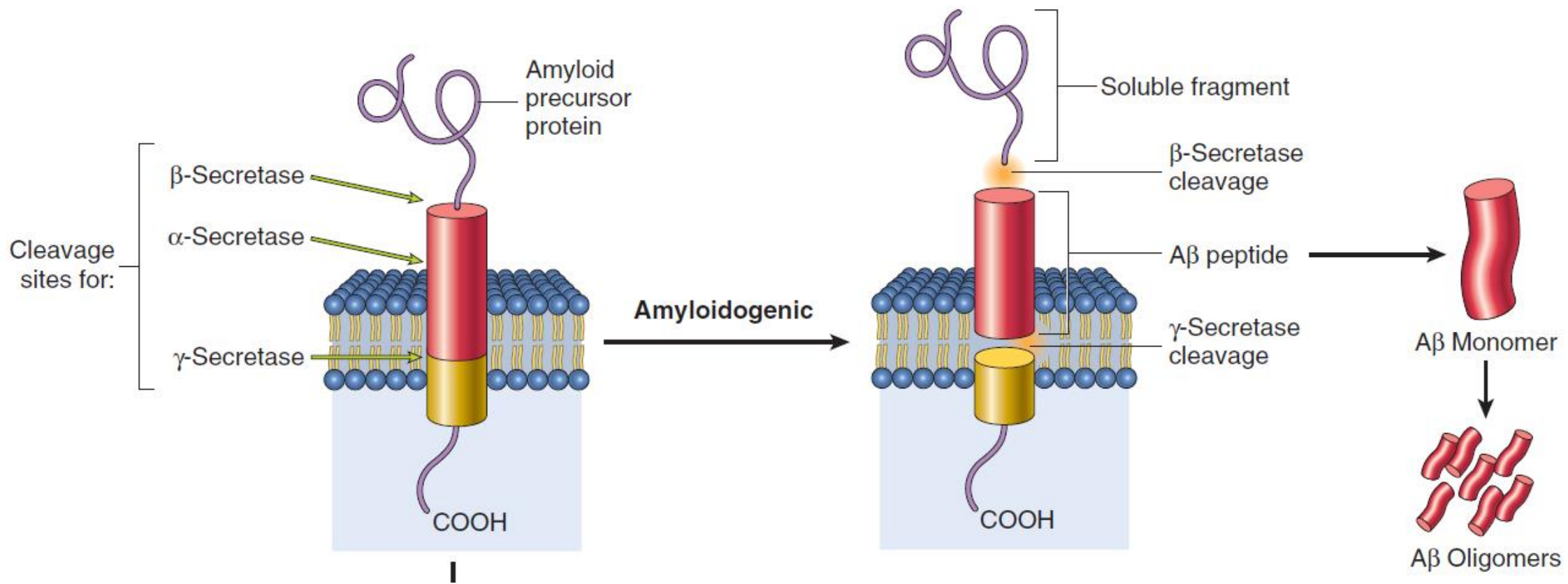
Role of A β

- ▶ AD results when the transmembrane protein (amyloid precursor protein APP) is sequentially cleaved by the **enzymes β -amyloid-converting enzyme (BACE) (B-secretase) and γ -secretase** creating A β .
- ▶ Normally, APP can be cleaved by **α -secretase and γ -secretase**, liberating a nonpathogenic peptide.
- ▶ Mutations in APP or in components of γ -secretase lead to familial AD.
- ▶ The *APP* gene is located on chromosome 21, increased risk in down syndrome
- ▶ Once generated, A β is highly prone to aggregation >>>> PLAQUES FORMATION >>> decreased number of synapses and alter their function >>> memory disruption.



Nonamyloidogenic

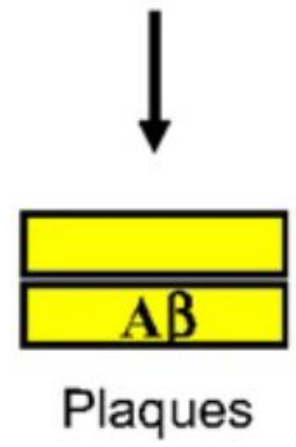
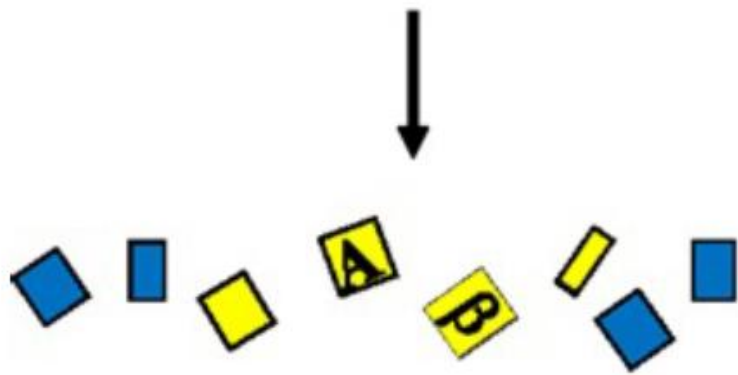


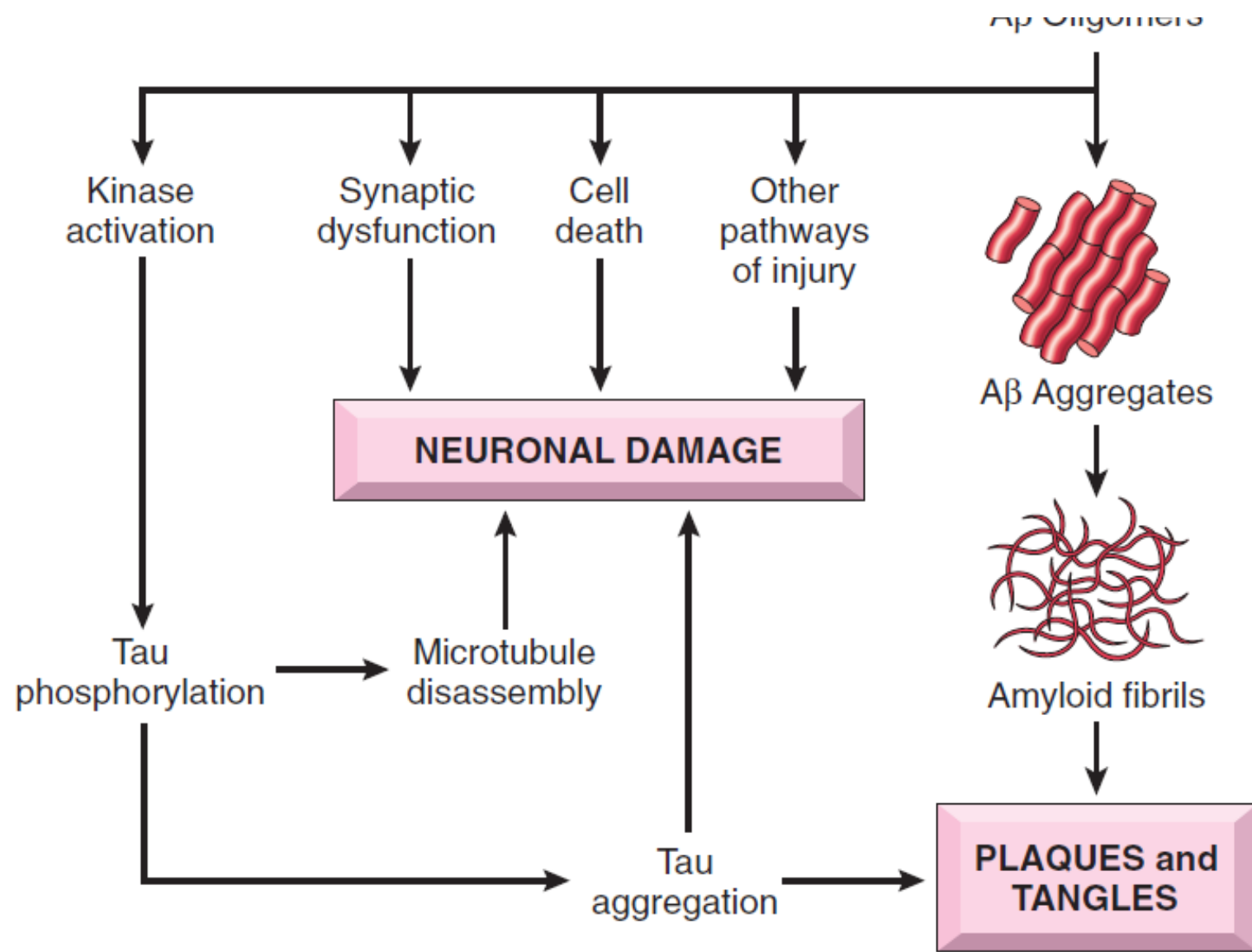




Normal

Amyloidogenic





Role of tau:

- ▶ Tau is a microtubule-associated protein.
- ▶ Present in axons in association with the microtubular network.
- ▶ Responsible for tangles in AD >>> Tau aggregates leads to cell death
- ▶ Hyperphosphorylated and loses the ability to bind to microtubules >>>> loss of microtubule stability >>> neuronal toxicity and death.

- ▶ Tau aggregates can be passed across synapses from one neuron to the next >>> spread of lesions.

Role of inflammation

- ▶ Innate immune system responds to A β and tau.
- ▶ Deposits of A β elicit an inflammatory response from microglia and astrocytes.
- ▶ Clearance of the aggregated peptide, and secretion of mediators that cause neuronal injury over time.

Basis for cognitive impairment

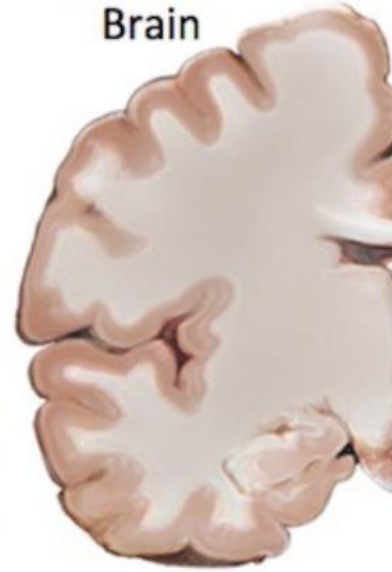
- ▶ Deposits of A β and tangles appear long before cognitive impairment
- ▶ In familial AD, deposition of A β and the formation of tangles precede cognitive impairment by as much as 15 to 20 years.
- ▶ Large burden of plaques and tangles is strongly associated with severe cognitive dysfunction.
- ▶ **The number of neurofibrillary tangles correlates better with the degree of dementia than does the number of neuritic plaques.**

Morphology

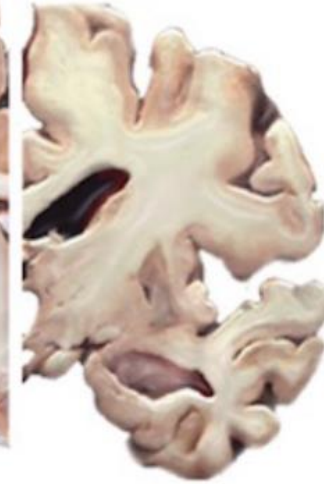
- ▶ Cortical atrophy,
- ▶ Widening of the cerebral sulci
- ▶ Most pronounced in the frontal, temporal, and parietal lobes.
- ▶ Compensatory ventricular enlargement (hydrocephalus ex vacuo).



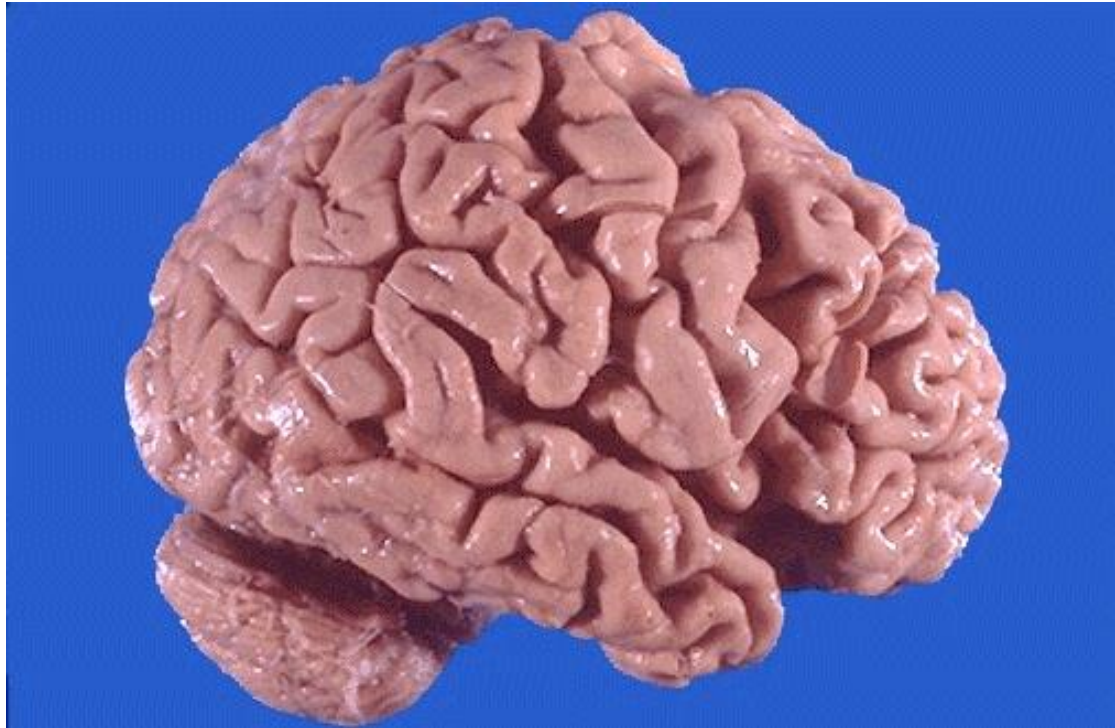
Healthy
Brain



Severe
Alzheimer's



Neuronal cell loss leading to extensive shrinkage in an Alzheimer's brain (right), as compared to a healthy human brain (left).



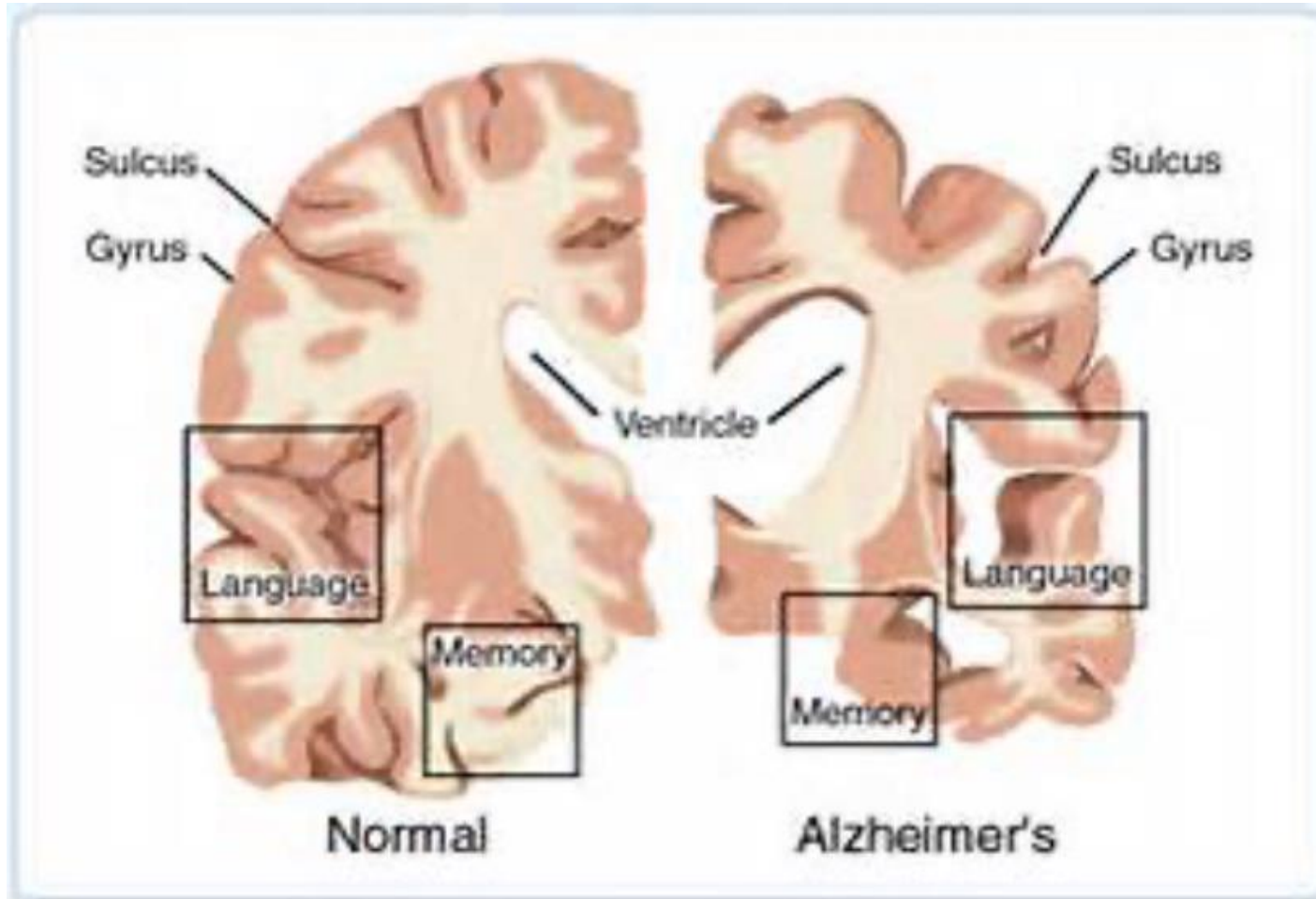
- ▶ Mainly in the frontal and parietal regions, characterized by **narrowed gyri** along with **widened sulci**.

- ▶ More marked atrophy seen superiorly and laterally, with sparing of the occipital region.



Progressive cortical atrophy with Alzheimer disease leads to compensatory dilation of the **cerebral ventricles** known as "hydrocephalus ex vacuo".



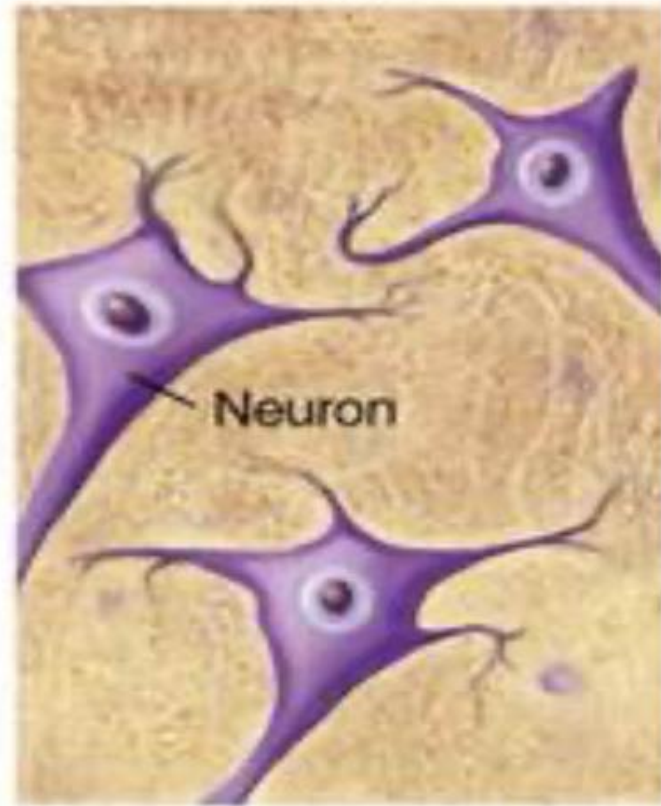


Alzheimer disease neuropathologic changes.

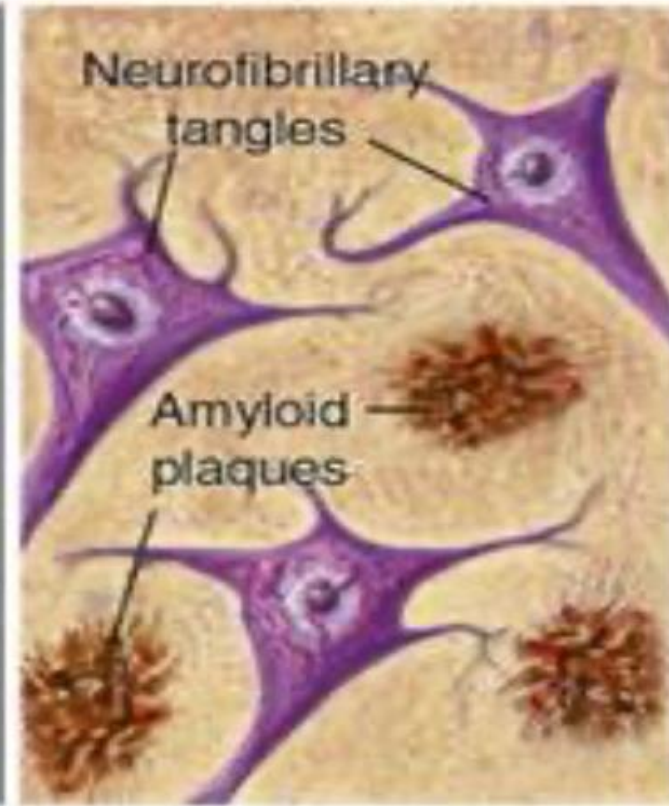
- ▶ **Neuritic plaques** (an extracellular lesion): central amyloid core surrounded by collections of dilated, tortuous, processes of dystrophic neurites.
- ▶ Hippocampus and amygdala and neocortex, (sparing of primary motor and sensory cortices until late)
- ▶ The amyloid core contains A β

- ▶ **Neurofibrillary tangles**, basophilic fibrillary structures in the cytoplasm of neurons, displace or encircle the nucleus; persist after neurons die, becoming extracellular.
- ▶ Cortical neurons, pyramidal cells of hippocampus, the amygdala, the basal forebrain, and the raphe nuclei.
- ▶ Hyperphosphorylated tau

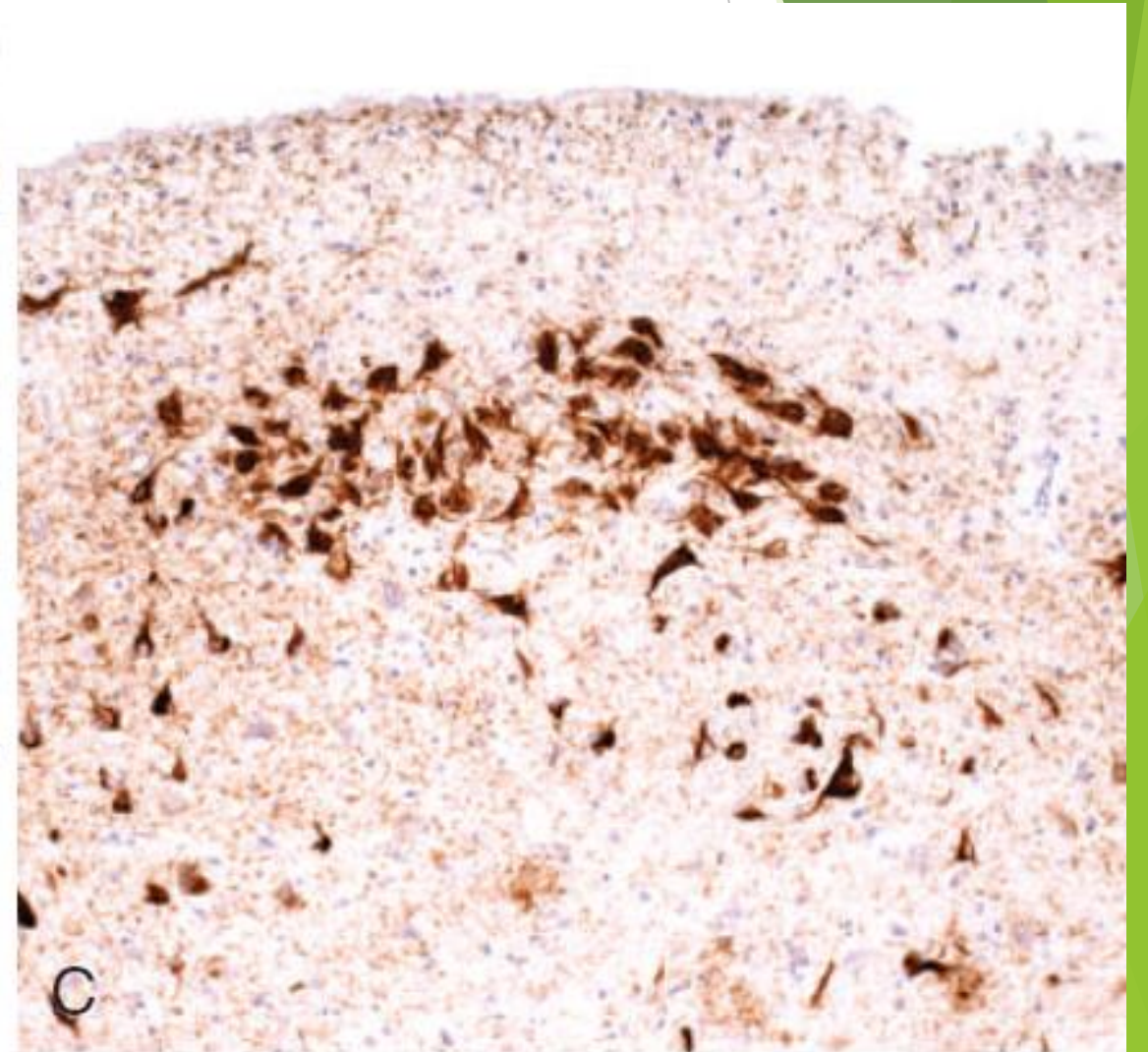
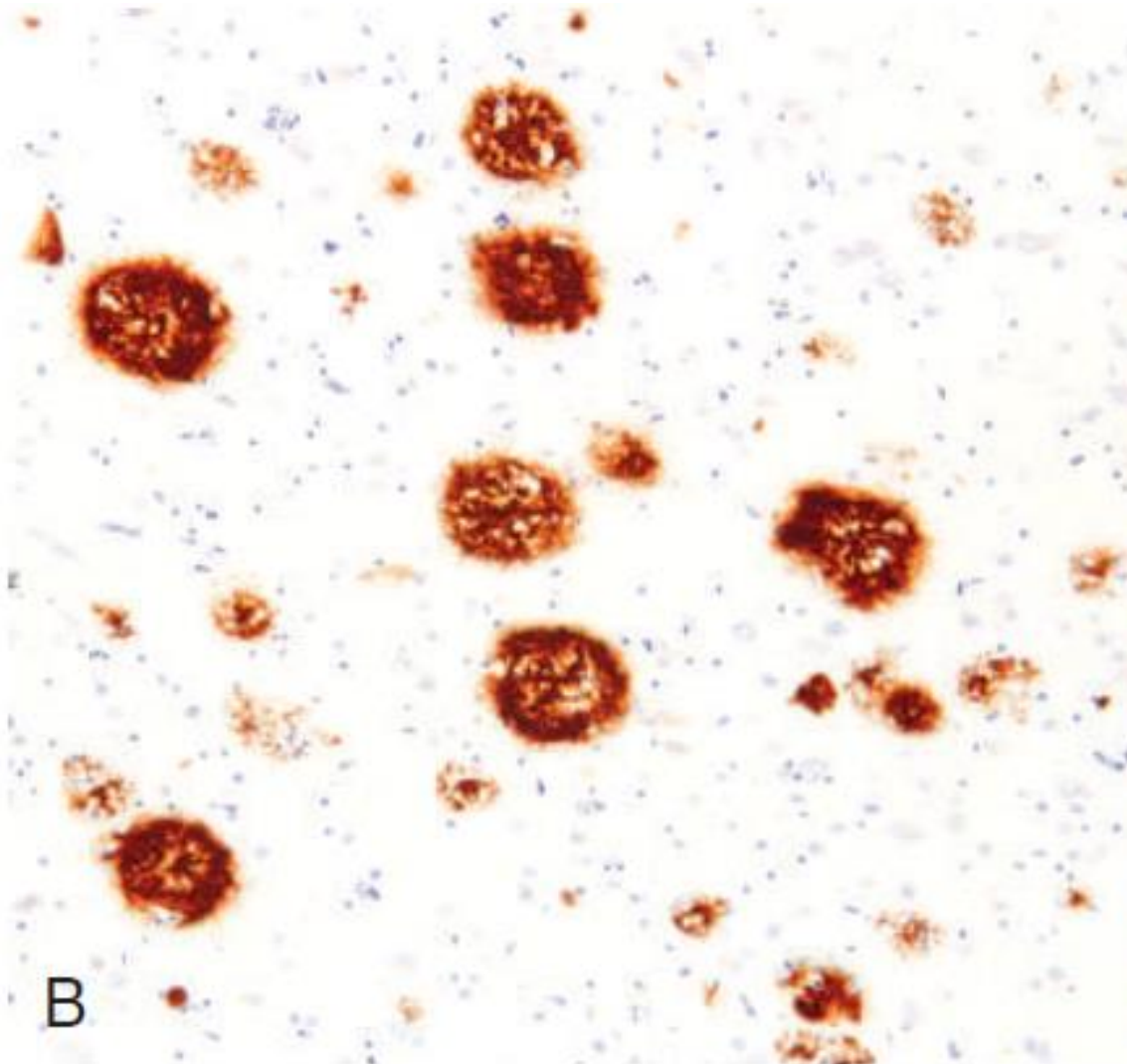
Normal

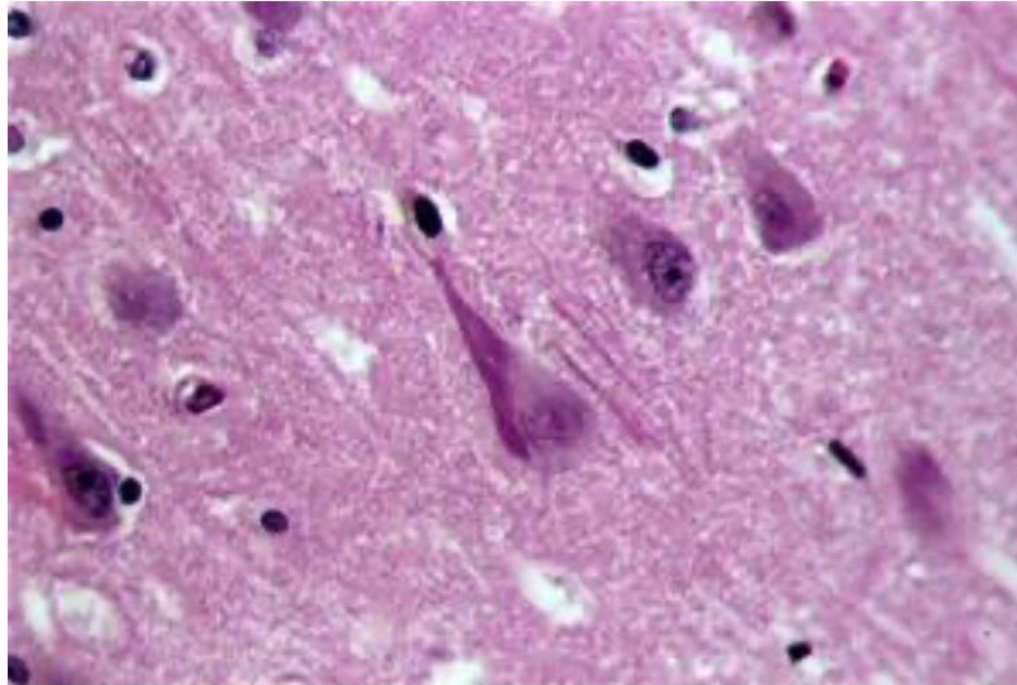


Alzheimer's

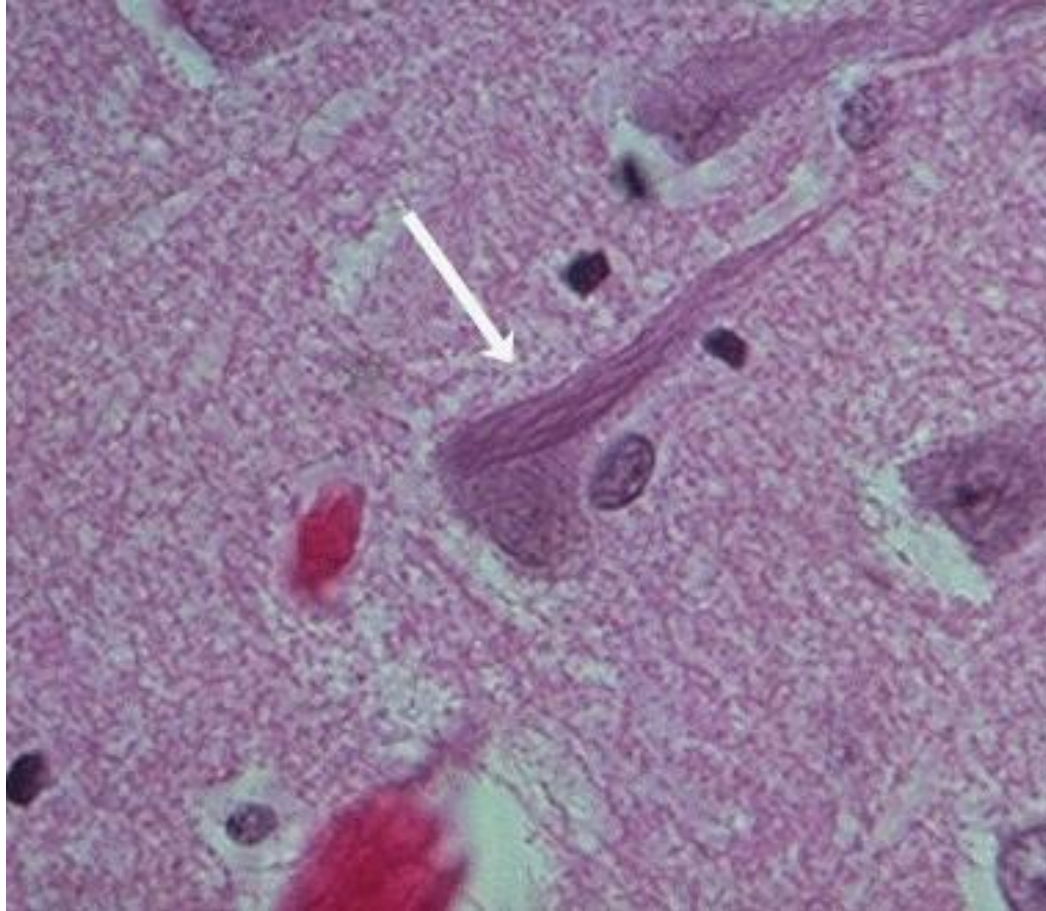


Plaques and tangles

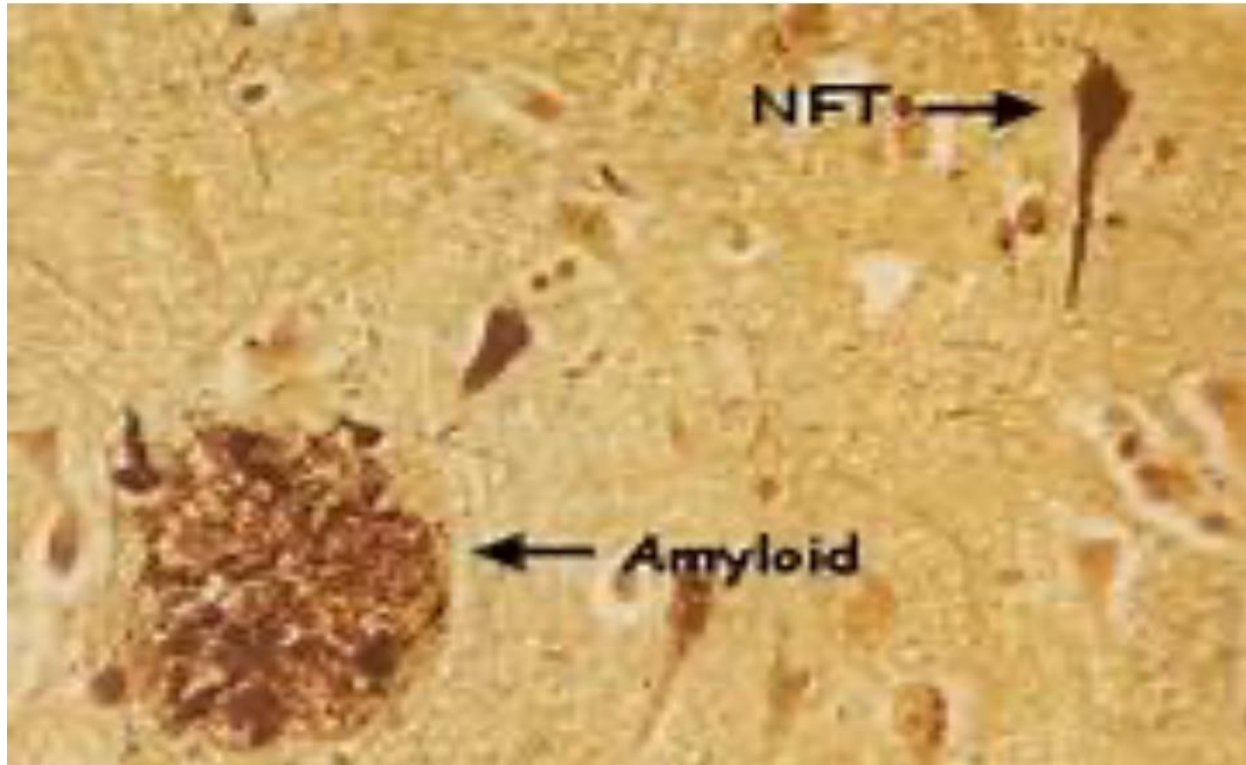


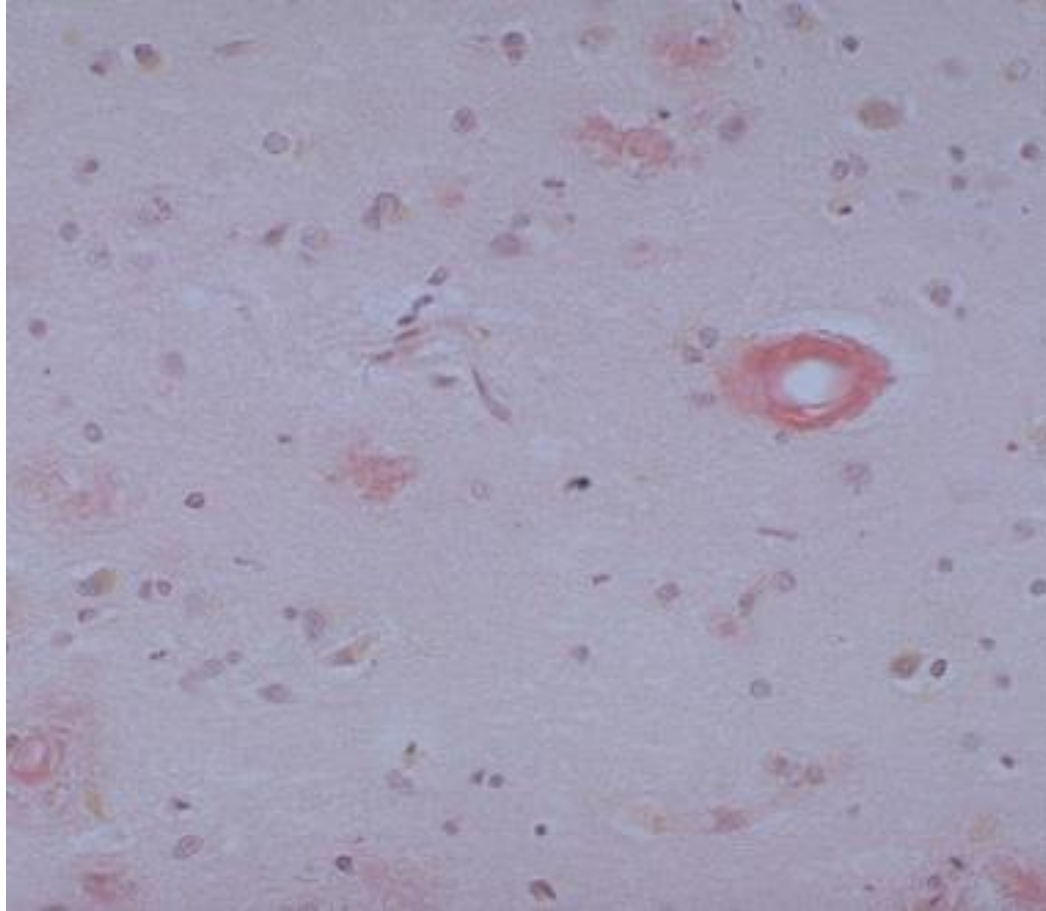


NEUROFIBRILLARY TANGLES

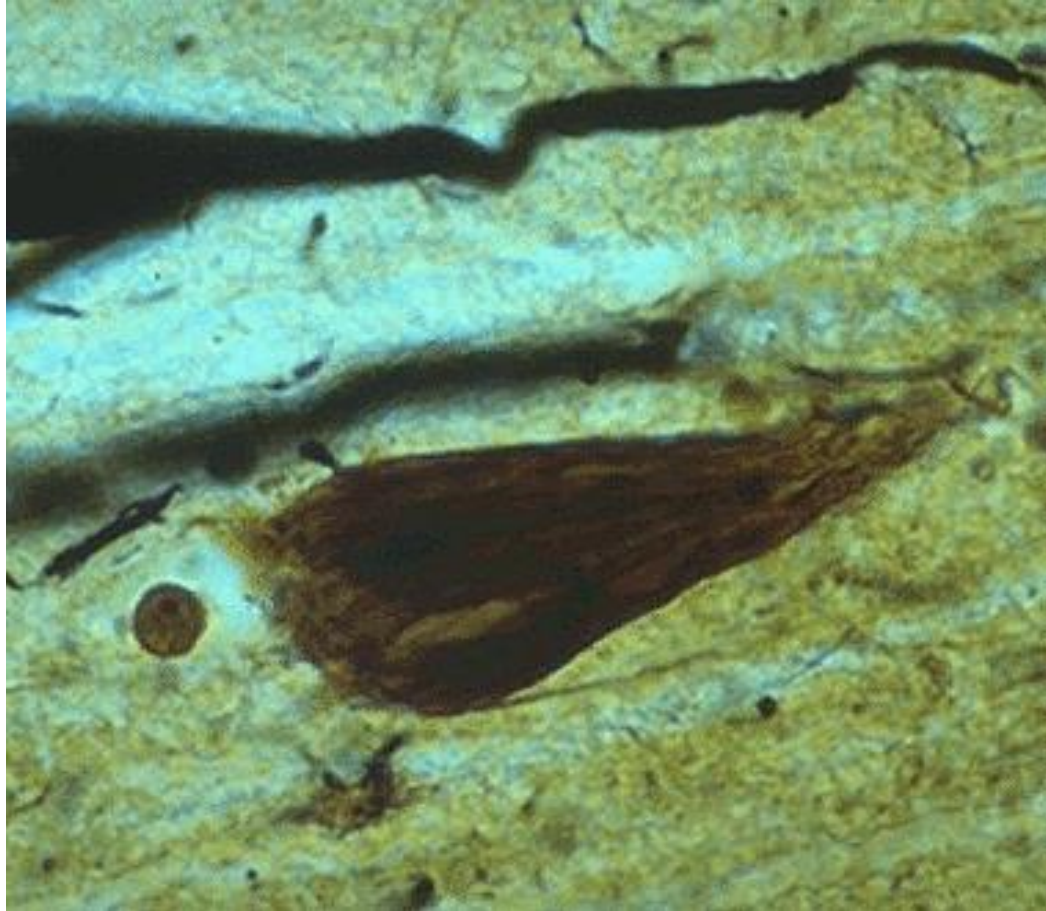


Neurofibrillary
tangles





Congo red
stain for
amyloid core
of plaques.

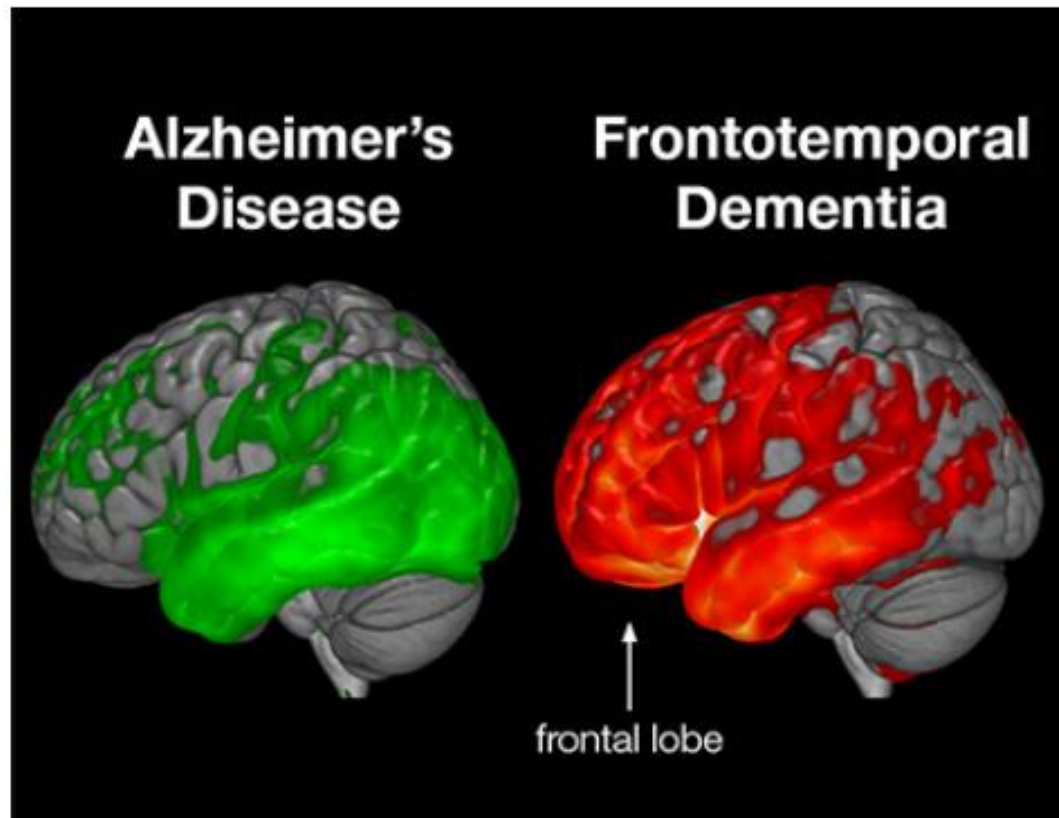


Silver stain for
NFT

Frontotemporal Lobar Degeneration

Frontotemporal dementias

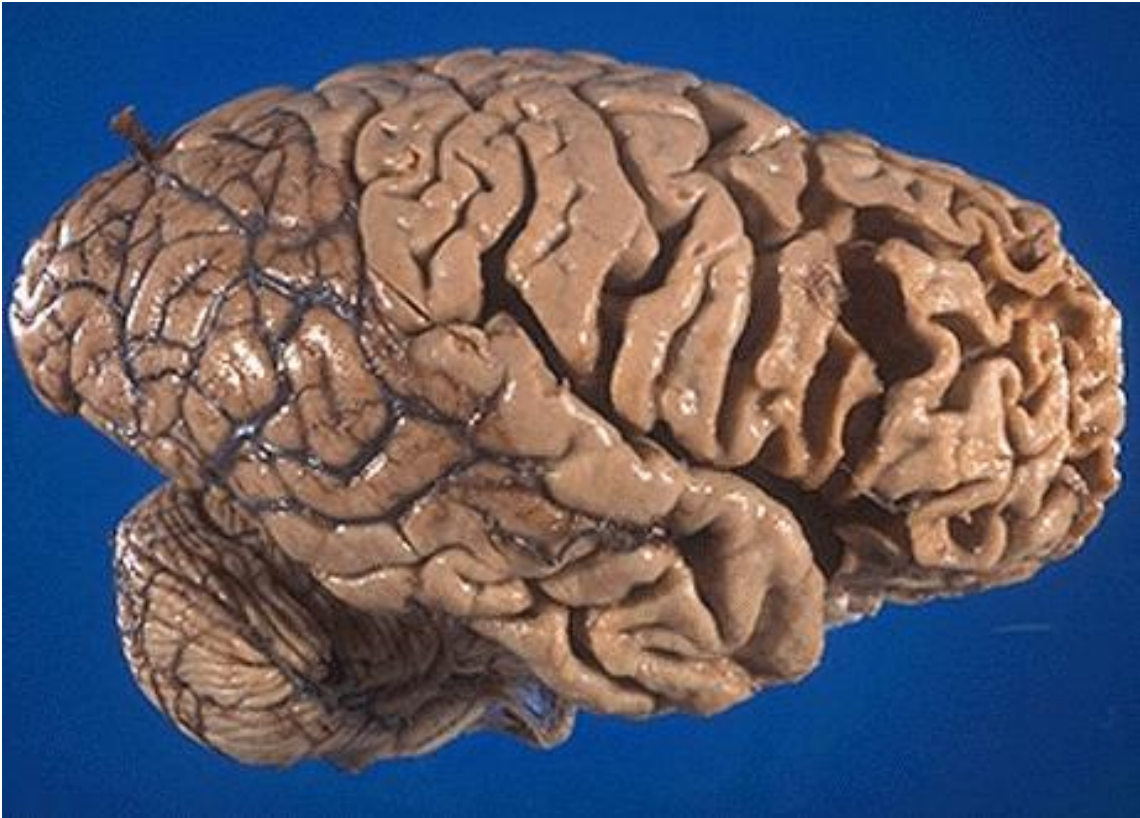
- ▶ Several disorders, preferentially affect **the frontal and/or temporal lobes**.
- ▶ Progressive deterioration of language and changes in personality
- ▶ **Behavioral and language problems precede memory disturbances, in contrast to AD.**
- ▶ The onset of symptoms occurs at younger ages than for AD.
- ▶ Neuronal inclusions, which may contain **tau or TDP43. (two forms of disease)**
- ▶ ***Pick disease*** (subtype of FTLD-tau), associated with smooth, round inclusions known as *Pick bodies*
- ▶ ***TDP34 subtype*** (also deposited in ALS)



- ▶ In AD there is sparing of the frontal lobe, at least at the beginning so behavioural changes are a late manifestation.
- ▶ In FTLD frontal is affected from the beginning so patients present with behavioural problems first.

MORPHOLOGY

- ▶ Atrophy of frontal and temporal lobes.
- ▶ Neuronal loss and gliosis
- ▶ In FTLD-tau, the characteristic neurofibrillary tangles, similar to AD
- ▶ Pick bodies in pick Disease.



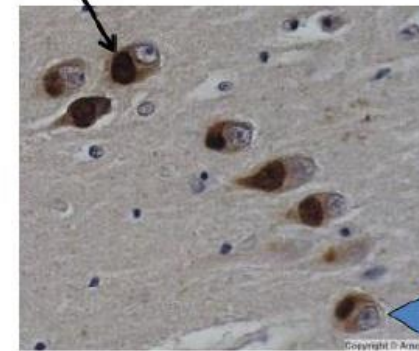
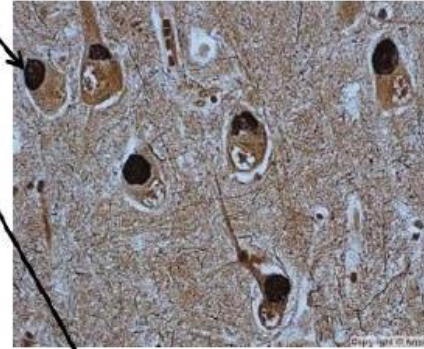
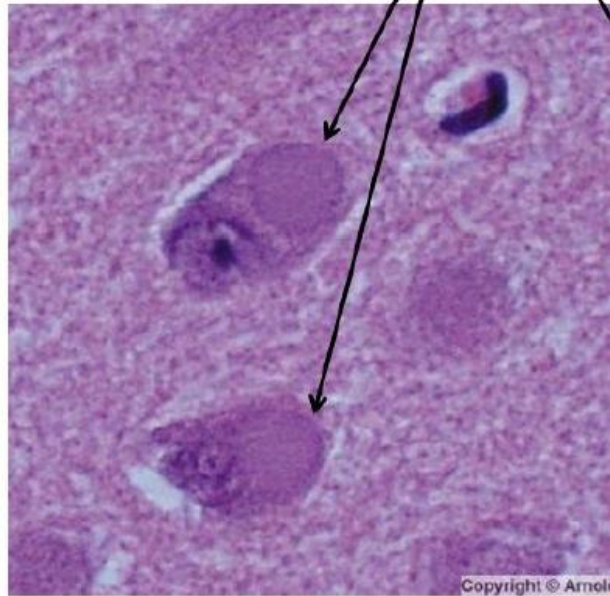
- ▶ Very marked **frontal lobe atrophy** and **temporal lobe atrophy**



Frontal lobes
are markedly
thinned

Pick bodies

Silver stain



Immunohistochemistry for Tau protein